Chapter 8 Sensory Transduction Network of *E. coli*

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Contents

Abstract The genome of *Escherichia coli* K12 encodes at least 6 classes of sensor proteins: 30 histidine protein kinases, 5 methyl-accepting chemotaxis proteins, 23 membrane components of the sugar:phosphotransferase system (PTS), 29 proteins with diguanylate cyclase and/or c-di-GMP-specific phosphodiesterase activity and two predicted serine/threonine protein kinases. The full signal transduction network additionally includes 32 response regulators, numerous chemotaxis proteins, PTS components, adenylate cyclase, CRP, and uncharacterized c-di-GMP-responsive components. Bacterial response to environmental signals can occur on several levels: the level of individual genes and proteins (changes in gene expression, posttranslational regulation), the whole-cell level (chemotaxis), and the multicellular level (biofilm formation). All signal transduction systems are energy-dependent but their energy expenditure is miniscule compared to that of the processes they

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regulate. A better understanding of the signal transduction mechanisms and integration of these mechanisms into the metabolic pathway model of the *E. coli* cell will remain major challenges for systems biology.

8.1 Introduction

For many years, *Escherichia coli* K12 served as a favorite model organism for studying principles and mechanisms of bacterial signal transduction. As a result, the current understanding of the signal transduction machinery in *E. coli*, albeit obviously incomplete, is probably as good as that for any organism in the prokaryotic or eukaryotic world. The availability of complete genome sequences of three strains of *E. coli* K12 (Blattner et al. 1997, Hayashi et al. 2006, Durfee et al. 2008) and their pathogenic counterparts (Hayashi et al. 2001, Perna et al. 2001, Welch et al. 2002, Johnson et al. 2007) made it possible to enumerate all (known) components of the signal transduction machinery encoded in each *E. coli* genome. This, in turn, allowed identification, at least in terms of sequence, of those signal transduction proteins whose biological functions are still unknown and remain to be experimentally characterized. In many respects, *E. coli* K12 proved to be a very convenient model: its signal transduction machinery is far more complex than that of its relatives who are obligate pathogens, such as *Haemophilus influenzae* or *Legionella pneumophila*. On the other hand, *E. coli* encodes far fewer signal transduction proteins than its free-living relatives (and opportunistic pathogens), such as *Pseudomonas aeruginosa*, *Shewanella oneidensis*, or *Vibrio cholerae*, not to mention the enormous expansion of signaling systems in the genomes of such model organisms as *Anabaena* PCC7120, *Myxococcus xanthus, or Streptomyces coelicolor* (Galperin 2005). Thus, signal transduction in *E. coli* is an experimentally tractable system that is responsible for much of the progress in understanding the principles and mechanisms of prokaryotic signal transduction.

The difficult task of a systematic description of the bacterial signal transduction machinery has been greatly simplified by the availability of specialized public databases, such as the *Mi*crobial *S*ignal *T* ransduction database (MiST, http:// genomics.ornl.gov/mist) at the Oak Ridge National Laboratory in Tennessee (Ulrich and Zhulin 2007) and the Kyoto Encyclopedia of Genes and Genomes (KEGG, http://www.genome.ad.jp/kegg/) at the Kyoto University in Japan (Kanehisa et al. 2008). The web pages of these databases dedicated to *E. coli* K12 (http://genomics. ornl.gov/mist/view organism.php?organism id=99, and http://www.genome.ad.jp/ dbget-bin/get pathway?org name=eco&mapno=02020, respectively) provide a bird's eye view of the composition and properties of signaling proteins encoded in the *E. coli* genome. In addition, the author maintains tables of Signal Transduction Census and Response Regulator Census at the web sites http://www.ncbi.nlm.nih. gov/Complete Genomes/SignalCensus.html and http://www.ncbi.nlm.nih.gov/ Complete Genomes/RRcensus.html, respectively. These web sites provide an easy way to access up-to-date information on signal transduction mechanisms in *E. coli* and related bacteria.

8.2 Diversity of Bacterial Signal Transduction Pathways

The two best-studied classes of membrane-bound receptor proteins are sensory histidine kinases and methyl-accepting chemotaxis proteins (MCPs), discovered in *E. coli* in the mid 1980s (Grebe and Stock 1999, Stock et al. 2000, Inouye and Dutta 2003). In the past several years, analyses of microbial genomes, as well as experimental studies, revealed several additional classes of bacterial receptors, which include Ser/Thr protein kinases and protein phosphatases, adenylate cyclases, diguanylate cyclases and c-di-GMP-specific phosphodiesterases (Table 8.1).

The signaling pathways utilized by various receptors are shown on Fig. 8.1 Signaling by histidine kinases and MCPs is usually referred to as two-component signal transduction, as it includes phosphoryl transfer between two different proteins, a histidine kinase and a response regulator. Two-component signal transduction pathways are extremely diverse but always include the following three steps:

Sensor type	N ₀ .	Function	Signaling mechanism
Histidine kinase	30	Transcriptional regulation, control of other processes	Phosphorylation of the REC domain of various response regulators
Methyl-accepting chemotaxis protein	5	Chemotaxis	Interaction with histidine kinase CheA, chemotaxis response regulator CheY
Ser/Thr protein kinase	$1 + 1^a$	Transcriptional regulation, posttranslational regulation	Phosphorylation of Ser or Thr residues in target proteins
Ser/Thr protein phosphatase	\overline{c}	Same as above	Dephosphorylation of Ser/Thr protein kinases or other target proteins
PTS membrane component	23	Sugar transport, chemotactic signaling, regulation of adenylate cyclase activity	Direct effect on chemotaxis, most likely through direct interaction of PTS enzyme I with the histidine kinase CheA
Adenylate cyclase	1	Global regulation of transcription	Synthesis of cAMP
Diguanylate cyclase	$12+7^b$	Regulation of protein and polysaccharide secretion	Synthesis of c-di-GMP
c-di-GMP-specific phosphodiesterase	$10+7^{\rm b}$	Same as above	Hydrolysis of c-di-GMP

Table 8.1 Principal Classes of Sensory Proteins in *Escherichia coli* K12

^a While YegI is believed to function as a Ser/Thr kinase, it remains unclear whether UbiB is an enzyme of ubiquinone biosynthesis or a Ser/Thr kinase that regulates this pathway (see the text for details).

^b Seven *E. coli* K12 proteins contain both GGDEF and EAL domains and could potentially catalyze both synthesis and hydrolysis of c-di-GMP (see the text for details).

Fig. 8.1 Signal transduction pathways of the principal classes of bacterial receptors. Signal transduction from a two-component signal transduction system (1), methyl-accepting chemotaxis sensor protein (2), phospho*enol*pyruvate-dependent sugar:phosphotransferase system (3), Ser/Thr protein kinase (4), adenylate cyclase (5), and sensor diguanylate cyclase (6)

(i) phosphorylation of a His residue in the kinase molecule; (ii) phosphoryl transfer to an Asp residue in the molecule of the cognate response regulator; (iii) conformational change of the response regulator that alters its interaction with its target on the chromosomal DNA or bacterial flagellum or, in some cases, the enzymatic activity of its output domain (see below).

Chemotaxis signaling, which starts from MCPs, is a special kind of twocomponent signal transduction that involves a specialized histidine kinase CheA, which directly interacts with MCPs, and a specialized response regulator CheY that consists of stand-alone receiver domain without any output domains. Regulation of flagellar motility is based on the interaction of the phosphorylated form of CheY with the FliM protein at the base of the flagellum, which affects the direction of flagellar rotation and thus regulates the chemotaxis response (Aizawa et al. 2002, Szurmant and Ordal 2004).

Components of the PEP-dependent sugar:phosphotransferase system (PTS) participate in phosphorylative sugar uptake and traditionally have not been considered part of the signal transduction machinery. Nevertheless, two members of the PTS phosphorelay play key roles in signal transduction. The phosphorylation level of the PTS enzyme I (EI) directly affects the chemotaxis machinery, whereas the phosphorylation level of the glucose-specific enzyme IIA ($EIIA^{Glc}$) modulates the activity of the adenylate cyclase, at least in *E. coli* and its closest relatives (Postma et al. 1993, Deutscher et al. 2006).

Ser/Thr protein kinases phosphorylate Ser and Thr residues in various cellular proteins. Only a small fraction of their targets have been identified so far. Ser/Thr protein phosphatases reverse the effect of Ser/Thr protein kinases by dephosphorylating their target proteins or, in some cases, the Ser/Thr protein kinases themselves (Shi et al. 1998, Deutscher and Saier 2005).

The adenylate cyclase modulates the cellular level of cyclic adenosine monophosphate (cAMP), a key cellular second messenger that regulates transcription from a variety of relatively weak promoters. The mechanism of this regulation includes binding of cAMP to a specialized adaptor protein, CAP (also referred to as cAMP receptor protein, CRP), triggering a conformational change in CRP that increases its affinity to DNA and allows it to activate transcription of otherwise poorly expressed genes (operons).

Signaling through diguanylate cyclases includes modulation of the cellular level of another cellular second messenger, cyclic dimeric bis-(3 –5)-guanosine monophosphate (c-di-GMP), which regulates a variety of function related to the cell surface elements, including motility, secretion of proteins and exopolysaccharides, biofilm formation, and production of certain virulence factors (Römling et al. 2005, Jenal and Malone 2006). Some of the c-di-GMP functions are mediated by its binding to the recently described PilZ domain, while others might involve other binding proteins, including diguanylate cyclases themselves. Cyclic-di-GMPspecific phosphodiesterases, which catalyze c-di-GMP hydrolysis, could also function as c-di-GMP-binding proteins.

8.3 Signal Transduction Machinery of *E. coli*

8.3.1 Two-component Sensors: Histidine Kinases

Histidine kinases are most numerous and most diverse membrane receptors encoded in bacterial genomes. Accordingly, they control the greatest variety of cellular responses. Most of the diversity of histidine kinases comes from the sensory (signal input) domains, which can be periplasmic, membrane-embedded or cytoplasmic. A single histidine kinase can contain several sensory domains, for example a periplasmic sensory domain and one or more ligand-binding PAS domains in the cytoplasm. In contrast, cytoplasmic signal transduction modules of histidine kinases are rather uniform and consist of two structural domains, dimerization/phosphorylation HisKA domain that consists of long alpha-helices and a C-terminal globular ATPase domain. Signal transmission by histidine kinases involved formation of dimers, so that an ATPase domain of one molecule binds ATP and transfers its γ -phosphate onto a conserved histidine residue in the HisKA domain of the other molecule in the dimer. This phosphoryl residue is subsequently transferred to an aspartyl residue in the receiver domain of the cognate response regulator. Analysis of sequence similarities between different histidine kinases by Parkinson and Kofoid (1992) revealed five conserved sequence motifs, referred to as H, N, G1, F and G2 boxes. The first of these boxes corresponded to the sequence motif around the conserved phosphorylaccepting histidine residue.

A cell of *E. coli* K12 encodes 30 histidine kinases; functions of six of them (AtoS, RstB, YehU, YpdA, YfhK, and YedV) still remain unknown (Hagiwara et al. 2004, Yamamoto et al. 2005), see Table 8.2 Among the remaining 24, by far the most (six, namely, BaeS, BasS, CpxA, EvgS, RcsC, and RscD), are involved in response to the envelope stress. Two more, EnvG and KdpD, are responsible for osmotic stress and adjustment of the magnitude of K^+ gradient. Other perceived signals include phosphate and/or its Ca^{2+} or Mg^{2+} salts (PhoQ, PhoR); nitrate and nitrite (NarQ, NarX); oxygen and/or hydrogen peroxide (ArcB, BarA); heavy metals, such as Cu^+/Ag^+ , (CusS) or Zn^{2+} and Pb²⁺ (ZraS); di- and tricarboxylates (CitA, DcuS); glucose-6-phosphate (UhpB), glutamine (GlnL), and trimethylamine N-oxide (TorS). One more histidine kinase sensor, QseC, is responsible for quorum sensing.

It is remarkable how many histidine kinases are sensing either envelope and osmotic stress or the redox state of the cell and the availability of terminal electron acceptors. The fact that these histidine kinases coexist in the same cell suggests a certain degree of sophistication in their interactions, seen, for example, in the complex division of functions between NarQ and NarX (Stewart 2003). In most cases, however, the hierarchy between different sensors, if any, remains unknown.

8.3.2 Two-component Transmitters: Response Regulators

Two-component response regulators are diverse proteins that share the common phosphoacceptor REC domain, often referred to as the CheY-like domain, after its best-known representative (Galperin 2006, Gao et al. 2007). This domain catalyzes phosphoryl transfer from the His residues of the histidine kinase HisKA domains to its own aspartate residues, as well as its own dephosphorylation (Thomas et al. 2008). The combination of these two activities in the REC domains of each particular response regulator determines the half-life of the phosphorylated form of

Histidine	Response	Signal	Regulated system or process
kinase	regulator		(genes)
ArcB ^a	$ArcA^b$	Redox state of the respiratory chain component(s)	Aerobic/anaerobic respiration
AtoS	A to Cd	Unknown (expression induced by acetoacetate)	Short-chain fatty acid metabolism (atoDAEB)
BaeS	BaeR ^b	Envelope stress	Multidrug efflux (mdtABCD, acrD)
BarA ^a	UvrY ^c	O_2 , H_2O_2 , oxidative stress	Carbon storage (csrB), catalase (katE)
BasS	BasR ^b	Envelope stress (high Fe^{2+})	Multidrug efflux
CheA	CheY, CheB	MCPs, PTS sugars	Chemotaxis
CitA (DpiB)	$CitBc$ (DpiA)	Citrate	Citrate metabolism (citCDEFG, citT)
CpxA	$CpxR^b$	Envelope stress, misfolded proteins	Protein degradation (htrA)
CreC (PhoM)	CreB ^b	Unknown (induced by growth in minimal media)	Central metabolism
CusS	CusR ^b	Cu^+ , Ag ⁺	Efflux transporters
DcuS	DcuR ^c	Fumarate, C4-dicarboxylates	Fumarate respira-tion (dcuB)
EnvZ	OmpR ^b	Envelope stress	Outer membrane (ompC, ompF)
EvgS ^a	EvgA ^c	Envelope stress	Multidrug efflux
GlnL (NtrB)	$GlnGd$ (NtrC)	Nitrogen starvation	Glutamine metabolism
KdpD	KdpEb	Osmotic stress	K^+ transport (kdpABC)
NarQ	NarP ^c	Nitrite/nitrate	Nitrate reductase (narGHIJ), formate dehydrogenase
NarX	NarL ^c	Nitrite/nitrate	Nitrate reductase (narGHIJ), formate dehydrogenase
PhoQ	PhoP _b	Low Mg^{2+}	Various genes
PhoR	PhoB ^b , PhoP	Low phosphate	Phosphate assimilation (phoA, phoB)
QseC	QseB ^b	Cell density (autoinducer-2), epinephrine, norepinephrine	Flagellar biosynthesis
RcsC ^a	RscB ^c	Unknown	Colanic acid biosynthesis
RscD	RscB ^c	Unknown	Colanic acid biosynthesis
RstB	RstA ^b	Unknown	Acid resistance, flagellar and capsular biosynthesis
TorS ^a	TorR ^b	Trimethylamine-N-oxide	TMAO reductase (torCAD)
U hp B	U hp A^c	UhpC, glucose-6-phosphate	Hexose phosphate uptake $(uhpT)$
ZraS (HydH)	ZraR ^d (HydG)	Heavy metals (Zn^{2+}/Pb^{2+})	Efflux transporter
YedV		Unknown	Unknown
YehU	YehT ^e	Unknown	Unknown
YfhK	YfhA ^d	Unknown	Unknown
YpdA	YpdB ^e	Unknown	Unknown
	FimZ ^c (YbcA)	Unknown	Fimbriae biosynthesis
	$RssB$ (Hnr)	Unknown	Proteolysis of RpoB by ClpXP

Table 8.2 Two-component signal transduction in *E. coli*

^a A hybrid histidine kinase that contains a receiver domain at its C-terminus.

^b DNA-binding transcriptional regulator, OmpR/PhoB (winged helix) family.

^c DNA-binding transcriptional regulator, NarL/FixJ (helix-turn-helix) family.

^d DNA-binding transcriptional regulator, NtrC (enhancer-binding) family.

^e DNA-binding transcriptional regulator, LytR/AgrA family.

the domain (CheY∼P or, more generally, REC∼P) and hence, the fraction of the response regulator molecules that are in the active (phosphorylated) conformation at any given time. A great majority of response regulators combine the REC domain with some kind of a signal output domain. However, some response regulators, such as the chemotaxis response regulator CheY, consist of a stand-alone REC domain. Chemotactic signal transduction through CheY relies solely on protein-protein interactions. Phosphorylation of CheY by phosphoryl transfer from the chemotaxis histidine kinase CheA shifts the CheY molecule into the active conformation that has an increased affinity to its target molecule FliM in the flagellar basal body. Non-phosphorylated CheY is also capable of interacting with FliM, albeit not as strongly. Thus, phosphorylation of CheY merely shifts the equilibrium of its two principal forms (there appear to be intermediate forms as well (Dyer and Dahlquist 2006)), leading to a change in the rotation pattern of the flagellum, which is reflected in an altered motility pattern of the whole cell.

With the exception of members of the CheY protein family, all other response regulators are two-domain (or three-domain) proteins that combine the REC domain with a signal output domain, which is usually located at the C-terminus of the polypeptide chain. Most of these proteins (in *E. coli*, 29 out of 32) are transcriptional regulators that activate or repress transcription of specific target genes. Accordingly, the most common output domains bind DNA, although some response regulators have enzymatic or ligand-binding output domains. The most common DNA-binding response regulators belong to the OmpR/PhoB family and have a winged helix-turnhelix DNA-binding domain. In *E. coli*, this family includes 14 proteins of the total of 32 response regulators (Table 8.2). The second in abundance with 9 representatives in *E. coli* is the NarL/FixJ family of response regulators with a typical helix-turnhelix DNA-binding output domain. Less common DNA-binding response regulators contain DNA-binding output domains of the Fis type (NtrC family) and LytTR type (LytR/AgrA family) with 4 and 2 representatives, respectively, encoded in the *E. coli* genome. Despite the differences in the structures of the DNA-binding response regulators, they all appear to follow a general mechanism of activation in response to the environmental signals. In each case, phosphorylation of the REC domain favors its transition into an active conformation and/or its dimerization (Toro-Roman et al. 2005, Gao et al. 2007). Dimerization of response regulators is a key mechanism of the transcriptional regulation by two-component systems, as response regulator dimers have a higher affinity to the tandem (or palindromic) transcriptional regulator binding sites on the chromosome. Within each family of response regulators, the signaling specificity is determined by the tight interaction of the REC domains with their cognate histidine kinases and of the HTH domains with the target sites on the DNA. As a result, transcriptional regulators with similar sequences (e.g., OmpR and PhoB) may have dramatically different biological functions. Some response regulators consist of more than two domains. In transcriptional regulators of the NtrC family (4 members in *E. coli*), the N-terminal REC domain and the C-terminal DNA-binding Fis-like domain are separated by the central AAA-type ATP-binding domain, whose ATPase activity is required for the DNA binding. In summary, bacterial response regulators contain a wide variety of output domains that put the

histidine kinases at the top of signaling hierarchy, allowing the cell to control its metabolism and behavior in response to various environmental challenges.

In some response regulators, the output domains are enzymatic. In *E. coli*, there is only one such response regulator, CheB, whose output domain is a methyl esterase of MCP proteins that takes part in chemotactic adaptation. Finally, *E. coli* and several closely related bacteria encode an unusual response regulator RssB (or Hnr), which regulates proteolysis of the stress sigma factor RpoS (Muffler et al. 1996, Zhou et al. 2001, Hengge-Aronis 2002). Its C-terminal domain is a degraded version of the Ser/Thr protein phosphatase domain which has apparently lost its catalytic activity and participates solely in protein-protein interactions (Galperin 2006).

8.3.3 Methyl-accepting Chemotaxis Proteins

Escherichia coli K12 encodes 5 methyl-accepting chemotaxis proteins (MCPs). The signals sensed by each of them have been experimentally characterized as follows: Tsr – serine; Tar – aspartate, maltose; Trg – ribose, galactose; Tap – dipeptides; and Aer – redox state of the respiratory chain (Szurmant and Ordal 2004). The last of these MCPs, Aer, is obviously important for sensing the presence of usable terminal electron acceptors, reflecting the choice between a respiratory and fermentative metabolism (Repik et al. 2000, Zhulin 2001). All these MCPs appear to interact with the chemotaxis histidine kinase CheA and transmit the respective signals thorough the two-component phosphorelay to the chemotactic response regulator CheY.

8.3.4 Phosphotransferase System Components

An MCP-independent mechanism of regulating chemotaxis is provided by the phospho*enol*pyruvate-dependent sugar:phosphotransferase system (PTS), which catalyzes uptake of certain sugars, coupling membrane transport of its substrates with their phosphorylation (Postma et al. 1993, Deutscher et al. 2006). Transport of sugar substrates by the PTS is coupled to signaling, both to the chemotaxis machinery and to the adenylate cyclase. Like histidine kinases, PTS proteins are phosphorylated on the histidine residue. However, in contrast to the ATP-His-Asp or ATP-His-Asp-His-Asp phosphorelay, typical for the two-component signaling, the PTS phosphorelay starts from phospho*enol*pyruvate (PEP) and includes only His residues, (at least, in EI, HPr and EIIA components). The high free energy of PEP hydrolysis ensures that in the absence of carbohydrate substrates all PTS components stay in the phosphorylated form. The limiting step in the whole phosphorelay appears to be PEP-dependent autophosphorylation of the first component, EI. Therefore, in the presence of carbohydrate substrates, phosphoryl flow through the PTS components occurs at a higher rate than re-phosphorylation of EI by PEP. As a result, EI, HPr and EIIA components become partly dephosphorylated, which serves as a signal both for the chemotaxis machinery and for the *E. coli* adenylate cyclase.

Although any direct interaction between PTS components and MCP or CheA remains to be demonstrated, the available data suggest that unphosphorylated EI can interact with CheA, modulating its activity and, hence, the cellular level of CheY∼P. The second mechanism of signal transduction from the PTS involves EIIAGlc. This protein has been shown to interact with the adenylate cyclase and other targets, including the lactose permease. The *E. coli* cell encodes 23 membrane components of the PTS, five of which (FruA, FrvB, FrwC, HrsA, and YpdG) are apparently specific to fructose. The other ones, according to the existing experimental data and sequence-based predictions, are specific to the following sugars: glucose (PtsG), mannose (ManX/Y), mannitol (MtlA, CmtA), N-acetylglucosamine (NagE), cellobiose (AscF, CelB), galactitol (GatC, SgcC), N-acetylgalactosamine (AgaC/D, AgaW), sorbitol (SrlA), maltose (MalX), trehalose (TreB), α -glucosides (GlvC), -glucosides (BglF), ascorbate (SgaB), and N-acetylmuramic acid (YfeV).

Thus, *E. coli* carries in its genome genes encoding chemotaxis receptors for almost any commonly found monosaccharide and several disaccharides. Whether these genes are constitutively expressed at sufficient levels to contribute to the cell behavior remains an open question. It appears that at least for some of the PTS receptor genes need to be induced by the corresponding sugar.

8.3.5 Ser/Thr Protein Kinases and Protein Phosphatases

Reversible protein phosphorylation on serine, threonine, or tyrosine residues is a key regulatory mechanism in eukaryotic cells. In the past several years, Ser/Thr protein kinases have been recognized in a variety of prokaryotic cells but are still often referred to as "eukaryotic-type" protein kinases. In certain groups of bacteria (e.g., actinobacteria) and archaea, Ser/Thr protein kinases appear to be the principal, if not the only (known) type of receptor proteins (Galperin 2005).

Most enterobacteria, including *E. coli*, encode just one or two Ser/Thr protein kinases and phosphatases, which remain poorly characterized. One of the predicted Ser/Thr protein kinases, UbiB, has been shown to be required for a hydroxylation step in ubiquinone biosynthesis and was initially thought to function as 2 octaprenylphenol hydroxylase (Poon et al. 2000). However, this enzymatic activity has not been experimentally demonstrated. In contrast, it has been identified as a member of the Ser/Thr protein kinase superfamily and has all the key active site residues intact. Thus, it remains unknown at this time whether UbiB is an enzyme of ubiquinone biosynthesis or a Ser/Thr protein kinase that regulates this process. The functions of the second predicted Ser/Thr protein kinase, YegI, also remain unknown.

8.3.6 Adenylate Cyclases

Bacteria encode several different variants (referred to as classes) of adenylate (adenylyl) cyclase, the enzyme that produces cAMP from ATP. The enzyme from *E. coli* is considered class I adenylate cyclase. It is a soluble enzyme that does not

appear to sense any environmental signals by itself. However, its activity is modulated by the EIA^{Glc} component of the glucose-specific phosphotransferase system. The phosphorylated form of EIIA^{Glc} appears to activate adenylate cyclase, whereas the dephosphorylated form, accumulating in the presence of extracellular glucose, does not bind to the adenylate cyclase or even inhibits it (Krin et al. 2002, Park et al. 2006). Thus, in the presence of glucose or other PTS sugars, adenylate cyclase activity decreases, leading to a drop in the cellular level of cAMP. This is one of the mechanisms contributing to the phenomenon of catabolite repression.

8.3.7 Diguanylate Cyclases and C-di-GMP Phosphodiesterases

A recently identified group of bacterial receptors includes proteins with so-called GGDEF and EAL domains that, respectively, synthesize and hydrolyze the second messenger c-di-GMP. Recent studies implicated c-di-GMP in regulating biofilm formation, development of flagellar apparatus, and a variety of other processes. The GGDEF domain has been shown to function as a diguanylate cyclase that produces a c-di-GMP molecule from two molecules of GTP (Paul et al. 2004, Ryjenkov et al. 2005). The EAL domain functions as c-di-GMP-specific phosphodiesterase, hydrolyzing c-di-GMP to a linear pGpG, and, eventually, to two molecules of GMP (Christen et al. 2005, Schmidt et al. 2005). *Escherichia coli* encodes 12 proteins with the GGDEF domain, 10 proteins with the EAL domain and 7 proteins that contain both of them and could potentially catalyze both reactions (Galperin et al. 2001, Galperin 2005). It appears, however, that in most of such fusion proteins, at least one of the domains is enzymatically inactive and serves to regulate the catalytic activity of the other one. In some cases, however, both domains appear to be active.

Our current knowledge of the functions of *E. coli* diguanylate cyclases and cdi-GMP-specific phosphodiesterases is very limited. The sensed ligand, oxygen (and/or CO and NO), has been established only for one of them, YddU, which was accordingly renamed 'direct oxygen sensor', or Dos (Delgado-Nixon et al. 2000). Several other GGDEF and EAL domain proteins, such as YaiC (AdrA), YdaM, YciR, and YhdA, have been shown to regulate, respectively, cellulose biosynthesis (Zogaj et al. 2001), production of curli fimbriae, and carbon storage, although the signal they respond to remains unknown. For other GGDEF and/or EAL domain proteins (Rtn, YcdT, YddV, YdeH, YeaI, YeaJ, YeaP, YedQ, YegE, YfeA, YfgF, YfiN, YhjK, YliF, YneF, YahA, YcgF, YcgG, YdiV, YhjH, YjcC, YlaB, YliE, YoaD), neither the sensed signal nor the regulated process are known at this time.

8.4 A System-level Look at the *E. coli* **Signal Transduction**

8.4.1 Multiple Responses to Multiple Signals

The above discussion shows that signal transduction machinery of *E. coli* is a complex network of interconnected pathways that underlie the ability of the cell to respond to environmental challenges. These responses are elicited by a variety

of environmental parameters and occur on several different levels, including the level of individual genes and operons (changes in gene expression), at the level of the whole cell (chemotaxis), and at the level of multicellular communication (quorum sensing, biofilm formation). The regulation of gene expression, in turn, is multi-faceted and can occur at the transcriptional level (changes in expression of certain genes, operons, or even global regulons), and at the levels of posttranscriptional (e.g. modulation of the mRNA decay rate) and post-translational regulation (e.g. modulation of enzyme activity, protein stability, or protein-protein interactions).

However, a closer look at the mechanisms actually utilized by *E. coli* shows that most of the cell responses occur either at the level of transcriptional regulation or at the level of whole-cell behavior (chemotaxis). The two-component signal transduction in *E. coli* is primarily targeted towards transcriptional regulation (29 of 32 response regulators are DNA-binding). Chemotaxis involves just two response regulators, CheY and CheB, and the single remaining response regulator (RssB or Hnr) acts post-translationally, at the level of proteolysis of RpoS (Hengge-Aronis 2002), and ultimately affecting transcription of RpoS-dependent genes. Another way transcription can be regulated by environmental signals is through the cAMP-CRP system. As mentioned above, sugar uptake by the PTS affects the adenylate cyclase activity and, hence, transcription from a variety of catabolite repressionsensitive promoters. Predicted Ser/Thr protein kinase YegI contains a C-terminal helix-hairpin-helix DNA-binding domain and is probably also involved in transcriptional regulation. There is a distinct possibility that transcription can also be regulated by signaling pathways leading from the cellular diguanylate cyclases. However, there is currently no experimental data to support that possibility.

The whole-cell behavioral changes include (i) chemotaxis in response to a variety of sugars, several amino acids, and/or changes in the redox state of the cell and (ii) production of exopolysaccharide and curli fimbriae, eventually leading to biofilm formation. This dichotomy might reflect the critical choice between "stay" and "run" survival modes, which appears to be governed by the c-di-GMP-mediated signaling.

The same cellular responses can be classified in terms of the environmental parameters that cause them. Although we still know very little about the signals sensed by several histidine kinases, predicted Ser/Thr protein kinase, diguanylate cyclases, and c-di-GMP-specific phosphodiesterases, the listing of the environmental parameters sensed by experimentally characterized histidine kinases, MCPs, and membrane components of the PTS shows two interesting trends. On one hand, the *E. coli* cell monitors (or, rather, is capable of monitoring) a variety of environmental stress conditions and extracellular concentrations of a variety of nutrients. The first group includes, among others, envelope stress, osmotic stress, presence of heavy metals, and presence of membrane-penetrating acids, such as acetate or benzoate. The second group includes a variety of hexoses, most disaccharides, di- and tricarboxylates, but apparently only one pentose (ribose) and only a minimal selection of amino acids (glutamine, serine, aspartate). While all these compounds are obviously important for *E. coli* metabolism, it is hard to rationalize why *E. coli* senses primarily hexoses

and not pentoses or these particular amino acids and not glutamate or asparagine. Some of these traits probably reflect simplification of the effector panel during adaptation of *E. coli* and other enterobacteria to the high-nutrient intestinal environment. Others might provide clues to the functional specialization of enteric bacteria within that specific ecological niche.

8.4.2 Energy Expenditure Considerations

It is important to note that environmental sensing is almost never energy-neutral: transmission of environmental signals requires significant energy expenditures, although minor in comparison to the energy requirements for motility, transcription of new genes (operons) or polysaccharide secretion,

As shown on Fig. 8.1, transmission of a signal through the two-component system takes an ATP molecule to phosphorylate a single molecule of a response regulator. Transcriptional regulation usually requires dimerization of response regulators, so two ATP molecules are being spent to convert an inactive response regulator into the active phosphorylated dimeric form. Autocatalytic spontaneous dephosphorylation of the receiver domains of response regulators weakens their protein-protein interaction, leading to the dissociation of dimers and a significant decrease in the DNA-binding ability. Therefore, the energy is spent here to achieve a rapid but relatively short-term activation (or, in some cases, repression) of transcription of certain genes (operons). Obviously, these energy expenditures are minor in comparison to the energy requirements of the transcription process, not to mention protein translation.

Transmission of the chemotactic signal includes a histidine kinase-response regulator pair and follows the same general principle as above. However, in case of CheB, as well as in the methylation-demethylation cycle, additional energy is being spent to regulate the adaptation time, i.e. to achieve a more precise timing of the signal. Again, these energy expenditures are minor in comparison to the energy spent on flagellar rotation that is required for motility of *E. coli*.

The lack of knowledge of the targets for Ser/Thr protein phosphorylation does not allow us to calculate the energy costs of this type of regulation. Nevertheless, they appear to be comparable to that of two-component signal transduction.

cAMP-mediated signaling requires a molecule of ATP to produce cAMP, which is then hydrolyzed to AMP by various phosphodiesterases. Converting the resulting AMP back to ATP requires two more ATP equivalents. Thus, transcriptional regulation of catabolite-sensitive operons requires at least 6 molecules of ATP per cAMP-CRP dimer.

In contrast, signal transmission through the PTS is remarkably energy efficient. As far as we know, chemotactic signaling by dephosphorylated EI and inhibition of the adenylate cyclase by dephosphorylated $EIIA^{Glc}$ do not require additional energy expenditure. However, the energy price here is paid in synthesizing all the components of the PTS and keeping them phosphorylated in the absence of the sugar substrate.

Finally, formation of a single c-di-GMP molecule consumes two molecules of GTP and four more ATP equivalents are required to restore these two molecules of GTP from pGpG. Further, the available crystal structures of c-di-GMP bound to proteins suggest that the active conformation of c-di-GMP is its dimer. The mechanisms of c-di-GMP-mediated regulation are still not fully understood, but both activation of cellulose biosynthesis through binding of c-di-GMP to the PilZ domain of the cellulose synthase (Amikam and Galperin 2005) and inhibition of flagellar formation through binding of the YcgR protein to the flagellar basal body (Ryjenkov et al. 2006) seem to occur solely by conformational changes, without any further energy-consuming reactions. Again, in these cases, energy expenditure seems to be minimal compared to that of the regulated process, that is, cellulose biosynthesis and export of flagellin.

In conclusion, despite the recent progress, there remain major puzzles in signal transduction pathways of even such well-studied organism as *Escherichia coli* K12. Determination of the range of signals sensed by this organism and the range of cellular responses elicited by these signals is an important goal of the ongoing experimental studies. A complete understanding of the signal transduction mechanisms and full integration of these mechanisms into the metabolic pathway model of the *E. coli* cell will probably remain a challenge for the nearest future.

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Chapter 9 Genome-Scale Reconstruction, Modeling, and Simulation of *E. coli's* **Metabolic Network**

Adam M. Feist, Ines Thiele, and Bernhard Ø. Palsson

Contents

Abstract Since the release of the first genome-scale metabolic reconstruction of the *E. coli* metabolic network in 2000, there has been a growing number of researchers around the world adapting it for a broad range of studies (Feist and Palsson 2008). The uses range from practical applications to obtaining basic biological understanding of cellular behavior. This range of uses is further expected to expand as the reconstruction broadens in scope and as new *in silico* methods are developed, implemented, and put to use.

In this chapter, we will describe foundational concepts central to the reconstruction process and model formulation, the history of reconstruction of the *E. coli* metabolic network, the development of reconstruction technology, genomescale constraint based modeling with key exemplary case studies of uses of the *E. coli* metabolic reconstruction, and insights into the future of the field. As such,

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this chapter should serve as a guide to those interested in either expanding the application of the *E. coli* reconstruction or adapting established applications to other organisms.

9.1 Foundational Concepts

The reconstruction of the *E. coli* metabolic network has led to the development of *'bottom-up'* reconstruction technology, genome-scale modeling methods, and basic and practical uses. A number of foundational concepts have also been developed during the period that we introduce here and provide background and a conceptual framework for the reader (see Palsson 2006, Price et al. 2004a).

Forming a BiGG knowledge base: A network reconstruction is based on a highly curated set of primary biological information for a particular organism; a biochemically, genetically and genomically structured (BiGG) knowledge base (Reed et al. 2006a). Such a knowledge base represents a large body of experimental data that is meticulously assembled and curated through the systems biology and reconstruction approaches detailed herein.

Genome-scale network reconstruction (GENRE): An organism-specific BiGG knowledge base is the basis for a GENRE. A GENRE is specific to a particular organism, for example, GENRE of *Escherichia coli* (below we will see four of these, specifically called *i*JE660, *i*JR904, *i*MBEL979, and *i*AF1260). A GENRE contains a list of all the known (and some predicted) chemical transformations that are believed to take place in the particular network (e.g. metabolic, transcriptional regulatory network, etc.).

The central role of network reconstruction in systems biology: Systems biology research generally can be conceptualized as a four-step process (Fig. 9.1). Foundational to the field is the generation of global, or genome-scale, data. The growing number of available 'omics' data types has created the need for formal and structured multi-'omic' data integration (Joyce and Palsson 2006). Omics data, along with legacy information (i.e., the 'bibliome') and detailed small-scale experiments, can be used to define the interactions among biological components that are used to reconstruct networks in particular organisms (Reed et al. 2006a). Network reconstruction is also an iterative, on-going process that continually integrates data in a formal fashion as it becomes available (Reed and Palsson 2003). These characteristics render the network reconstruction as a common denominator for those studying systems biology. The reconstruction effectively represents a 2-D annotation of a genome detailing not only the parts for an organism, but the interactions between specific components (Palsson 2004). Genome-scale reconstruction technologies for metabolic (Reed et al. 2006a), transcriptional regulation (Covert et al. 2004, Gianchandani et al. 2006, Herrgard et al. 2004) and signaling networks (Papin et al. 2005) have been established, and transcriptional/translational network reconstruction methods are currently under development (Thiele et al. 2009). An in depth review on the bottom-up reconstruction process (Palsson 2006) as well as a current review of biological network reconstruction (Feist et al. 2009) have been generated.

Fig. 9.1 Systems Biology as a 4-step Process. Step 1, the process is based on a variety of highthroughput data sets (i.e., 'omics' data) and a comprehensive assessment of the literature (i.e., bibliomic data). Step 2, all of the data types are used to reconstruct the list of biochemical transformations that make up a network as well as their genetic basis (Reed et al. 2006a). In principal, the network is unique. Step 3, the data contained in the reconstruction can be formally represented (i.e., in the form of matrices and logical statements) that can be mathematically characterized by a variety of methods. Step 4, the computational model enables a broad spectrum of applications, as reviewed in this chapter. Figure adapted from (Feist and Palsson 2008, Palsson 2006)

Constraint-based reconstruction and analysis (COBRA): COBRA is the overall philosophy and approach of applying constraints to limit the range of achievable functional (phenotypic) states of GENREs (outlined below). A GENRE operates under defined constraints. These constraints fall into at least four categories (Palsson 2006): physico-chemical, topological, regulatory, and environmental. Such constraints can be mathematically represented and imposed on the functional states that a GENRE can take on. Functional states can be assessed using a variety of computational methods (Palsson 2006, Price et al. 2004a) and have been disseminated in the form of a COBRA Toolbox (Becker et al. 2007) that is a MATLAB (The MathWorks Inc., Natick, MA) based software package.

Converting network reconstructions into a Genome-scale Model (GEM): A GENRE can be converted into a mathematical form (i.e., an *in silico* model) and used to computationally assess phenotypic properties (reviewed in (Price et al. 2004a)). The COBRA approach is used to analyze the properties of GENREs by assessing allowable functional states. Genome-scale reconstructions are thus a key step in quantifying the genotype-phenotype relationship and can be used to 'bring genomes to life' (Frazier et al. 2003). The availability of reconstructed metabolic networks for microorganisms has increased rapidly in recent years and a growing number of research groups are synthesizing GENREs for target organisms of interest (see Fig. 9.4) (Feist et al. 2009, Reed et al. 2006a).

The conversion of a reconstruction (GENRE) to an *in silico* model (GEM), represented by the arrow from step 2 to step 3 in Fig. 9.1, involves a subtle, but critical, transition. The chemical transformations of which a GENRE is comprised can be represented stoichiometrically (as well as other formats, e.g., a directed graph). Stoichiometric representations form a matrix, the rows of which represent the compounds, the columns of which represent the chemical transformations, and the entries of which are the stoichiometric coefficients (see section below and Fig. 9.6) With the definition of systems boundaries and other details, a network reconstruction can be converted into a mathematical format that can be computationally interrogated. The process that this arrow represents is the bridge between the realms of high-throughput data/bioinformatics and systems science.

9.2 History of the *E. coli* **Metabolic Network Reconstruction: An Ongoing and Iterative Process**

The 18-year history of metabolic reconstruction for *E. coli* is outlined in Fig. 9.2 (Feist and Palsson 2008, Reed and Palsson 2003). *E. coli* served as a model organism in the era of discovery of metabolic biochemistry, and thus, comprehensive metabolic reconstructions were developed before its genome sequence was available (Varma et al. 1993a,b).With the publication of the *E. coli* genomic sequence in 1997 (Blattner et al. 1997), the development and use of the metabolic reconstruction in *E. coli* grew rapidly in scope.

Fig. 9.2 The ongoing reconstruction of the E. coli metabolic network. History of the *E. coli* metabolic reconstruction. Shown are six milestone efforts contributing to the reconstruction of the *E. coli* metabolic network. For each of the six reconstructions (Edwards and Palsson 2000, Feist et al. 2007, Majewski and Domach 1990, Pramanik and Keasling 1997, 1998, Reed et al. 2003, Varma et al. 1993a,b) (see text for details), the number of included reactions (*diamonds*), genes (*triangles*), and metabolites (*squares*) are displayed. Also listed is the expansion in scope in each successive reconstruction. The start of the genome era in 1997 (Blattner et al. 1997) marked a significant increase in scope. The reaction, gene, and metabolite values for pre-genomic era reconstructions were estimated from the content outlined in each publication and in some cases, encoding genes for reactions were unclear. Fig. adapted from (Feist and Palsson 2008)

Pre-genome era: Beginning in 1990, a network reconstruction consisting of 14 reactions (characterizing primarily the TCA cycle and partially glycolysis) was generated to analyze the production and secretion of acetate during aerobic growth on glucose (Majewski and Domach 1990). This example demonstrates the scope of initial uses of network reconstructions of *E. coli*. Later, in 1993, a larger metabolic reconstruction consisting of 146 reactions was generated, representing key catabolic and anabolic metabolic pathways (Varma et al. 1993a,b). This reconstruction was used for computing (Varma et al. 1993a, Varma and Palsson 1993, 1994, 1995):Optimal production of cofactors and biosynthetic precursors, Maximum allowable generation of amino acids and nucleic acids, and Internal network flux distributions for optimal and sub-optimal growth.

The computational predictions based on the model were compared to experimental data and found to be consistent with measurements under both aerobic and anaerobic glucose minimal media conditions (Varma and Palsson 1994). The comparison of computation and experimental findings in this work demonstrated the important concept of comparison to *in vivo* data as computational outcomes have to be considered as hypotheses that need experimental confirmation.

Following these developments in the early 1990s, an expanded reconstruction consisting of 317 reactions was generated in 1997. It included cofactor and cell wall biosynthesis, and other additional metabolic pathways (Pramanik and Keasling 1997, 1998). This expanded reconstruction was used for computations that incorporated measured metabolite uptake and secretion rates to predict central metabolic fluxes which were found to be consistent with enzymatic flux values determined from isotopomer-based measurements (Pramanik and Keasling 1997, 1998). These studies also incorporated a growth rate dependent biomass objective function that had not been considered in previous studies. It should be noted that isotopomer-based measurements are also network dependent and studies are currently emerging looking specifically at this issue (Suthers et al. 2007).

Note that these pre-genome era reconstructions of *E. coli* metabolism were based solely on biochemical information and provided an important foundation for subsequent work at the genomic scale.

Genome era: The complete genome sequence for *E. coli* K-12 MG1655 was published in 1997 (Blattner et al. 1997). Its availability fueled a significant increase in network reconstruction content and scope as the genome sequence directly provided a list of parts (components) present in *E. coli* (Fig. 9.2). Utilizing the annotated sequence, a genome-scale metabolic reconstruction was generated for *E. coli* consisting of 627 unique reactions catalyzed by 660 gene products (Edwards and Palsson 2000). This reconstruction, later titled *i*JE660, was initially used to:Predict the phenotypes for knock-out mutants of the central metabolic pathways (Edwards and Palsson 2000), Design quantitative experiments (Edwards et al. 2001), and Predict the outcome of adaptive evolution in the context of the metabolic machinery available to the cell (Ibarra et al. 2002). These results demonstrated the utility of the reconstruction to understand growth characteristics of *E. coli*, the effects of gene deletions, and to point to areas of computational and experimental disagreement that identify targets for further biochemical characterization (see below).

An updated annotation of the *E. coli* K-12 MG1655 genome (Serres et al. 2001) and continual functional characterization of *E. coli* metabolic content enabled an expansion of the reconstruction in 2003, which consisted of 931 reactions catalyzed by 904 gene products (Reed et al. 2003). This reconstruction, titled *i*JR904, was an improvement over previous efforts in that contained both charge and elemental balancing of all reactions, expanded the various carbon source utilization pathways, contained a larger number of characterized transport systems and their encoding genes, better accounted for quinone usage in the electron transport chain, and better detailed the relationship between given genes, proteins, and reactions contained in the reconstruction (the GPR associations).

This reconstruction has been utilized for a broad number of applications reviewed later in this chapter. Utilizing the *i*JR904 (Reed et al. 2003) reconstruction, an expanded reconstruction of *E. coli* was generated (containing 979 reactions and titled *i*MBEL979) for the purpose of designing overproducing strains in the software framework MetaFluxNet (Lee et al. 2005).

The most recent metabolic reconstruction for *E. coli*, titled *i*AF1260, incorporates data from the most recent *E. coli* K-12 MG1655 genome annotation (Riley et al. 2006) and consists of 2,077 reactions and 1,260 genes (Feist et al. 2007). The advancements represented by *i*AF1260 over *i*JR904 lie in five main areas: an increased scope with the inclusion of 357 additional ORFs; compartmentalization into three distinct compartments (cytoplasmic, periplasmic and extra-cellular); the detailing of all grouped, or lumped, reactions (most often associated with lipid and lipopolysaccharide biosynthesis); the incorporation of reaction thermodynamics, calculated Gibbs free energy (Δ G°) values for 950 metabolites and 1935 reactions; and alignment with the EcoCyc database (Keseler et al. 2005) which provided expanded coverage for the network and content mappings for further computational analyses.

This 18-year history of reconstruction of the *E. coli* metabolic network has culminated in a network containing a total number of 1,260 metabolic genes covering 28% of the 4,453 identified ORFs on the *E. coli* genome. More importantly, the 1260 ORFs represent 48% of the functionally annotated ORFs that have been confirmed by experimental data (Table 9.1). Thus, 92% of the 1,260 gene products included in *i*AF1260 have been experimentally verified (Riley et al. 2006) with the balance of 8% having a computationally predicted function which necessitate confirmation with focused experimentation. Model-aided gap-filling and discovery will aid in this process (see Section 9.5.2). In addition, protein structures (computed or experimental) are available for a large fraction of the proteins in *i*AF1260 (Berman et al. 2000). Integration of protein structural data with the functional content of the reconstruction will lead to a better understanding of structural motifs and their properties.

Reconstruction of the *E. coli* metabolic network is thus approaching exhaustion of known metabolic gene functions and is now being used in a prospective fashion to discover new metabolic capabilities in *E. coli* (see below). As a result of this endeavour, the reconstruction of the *E. coli* metabolic network represents the bestdeveloped genome-scale network to date.

^a tabulated on a reaction basis, not counting outer membrane non-specific porin transport.

^b reactions can occur in or between multiple compartments and metabolites can be present in more than one compartment.

^c diffusion reactions do not include facilitated diffusion reactions and are not included in this total if they can also be catalyzed by a gene product at a higher rate.

^d overall genome coverage based on 4453 total ORFs in *E. coli*; *i*AF1260 contains 48% of the ORFs in *E. coli* that have been characterized experimentally (2403 ORFs).

9.3 Continuing Development of Reconstruction Technology

Development of the reconstruction process for metabolic networks: As illustrated in the previous section, the reconstruction process for metabolic networks is an iterative procedure that requires different types of experimental data and techniques at each phase of reconstruction. The experience with *E. coli* has led to

Fig. 9.3 The phases and tools necessary to generate a metabolic reconstruction. The genome-scale metabolic reconstruction process can be broken down into four major phases (center column), with each of the latter phases building off the previous. This process is iterative and driven by experimental data (primarily in the three latter phases). For each phase, specific data types are necessary and these range from high-throughput data types (e.g., phenomics, metabolomics, etc.), to detailed studies characterizing individual components (e.g., biochemical data for a particular reaction). For example, the genome annotation can provide a parts list of a cell, whereas genetic data can provide information about the contribution of each gene product towards a phenotype (e.g., when removed or mutated). The product generated from each reconstruction phase can be utilized and applied to examine a growing number of questions with the final product having the broadest applications

the formulation of the workflows that underlie metabolic reconstruction. The four phases of the reconstruction process are depicted in Fig. 9.3 and the product at each phase can be used for different applications, with the number of applications increasing with network development. This procedure represents the current status of network reconstruction, and the most recent *E. coli* reconstruction, *i*AF1260, was built accordingly (Feist et al. 2007) with the advantage of starting from an already well-established reconstruction, *i*JR904. The end product of this reconstruction effort is a platform for design and discovery, and key examples of use are given later in this chapter. More extensive descriptions exist, which outline the conceptual basis (Reed et al. 2006a) and the detailed process to generate genome-scale biological networks, (Feist et al. 2009) and these will not be repeated here.

Development of the reconstruction process beyond metabolism: The development and use of genome-scale reconstruction was rapid and many computational models were developed to address a growing spectrum of basic research and applied problems. Still, further development of reconstruction technology is necessary. The scope of reconstructions is bound to grow, representing more and more BiGG knowledge in the structured format of GEMs (Breitling et al. 2008). Growth in scope is likely to proceed in phases (Feist and Palsson 2008). Growth in scope in the nearterm will involve the transcriptional and translational machinery (Allen and Palsson 2003, Mehra and Hatzimanikatis 2006, Thiele et al. 2009, Thomas et al. 2007). Such an extension will enable a range of studies including the direct inclusion of proteomic data, fine graining of growth requirements, and the explicit consideration of secreted protein products.

Another expansion in scope in the near-term is the reconstruction of the genomescale transcriptional regulatory network (TRN). Such reconstruction at the genomescale is now enabled by new experimental technologies, such as ChIP-chip (Lee et al. 2002). Experimental interrogation of the currently available TRN suggests that we know about one-fourth to one-third of its content (Covert et al. 2004), indicating that there is much to be discovered. This expectation is being confirmed with highresolution ChIP-Chip data for *E. coli* (Cho et al. 2008). Once reconstructed, the TRN will allow computational predictions of the context-specific uses of the *E. coli* genome and the responses of two-component signaling systems.

Mid-term expansions in scope are likely to include the growth cycle, shock responses (e.g. heat and acid shock), and additional cellular functions (e.g. DNA replication and flagellar biosynthesis). Such a reconstruction should eventually be a comprehensive representation of the chemical reactions and transformations enabled by *E. coli*'s gene products.

Longer-term reconstruction may begin to address the 3-dimensional organization of the bacterial cell. In particular, high-resolution ChIP-chip data on the DNA binding protein could enable the estimation of the topological arrangement of the genome, and potentially elucidate the structure of the cell wall and other cellular structures that will allow a full 3-dimensional reconstruction of *E. coli*.

The two near-term expansions in content will encompass the activity of approximately 2000 ORFs in the *E. coli* genome. Clearly, quality-controlled reconstructions will help in guiding us to comprehensive genome-scale representation of all major cellular processes in bacteria at the BiGG data level of resolution that, in turn, enables GEMs of growing coverage and resolution. The scope of this effort has been described as being; "... 10 times more ambitious and 100 times more important for mankind [compared with Human Genome Project]..." Hans Westerhoff (Holden 2002).

Influence of the *E. coli* **reconstruction on the** *in silico* **analysis of other micro-organisms**: The metabolic network reconstruction of *E. coli* has been influential in the generation of other organism-specific metabolic networks. The *E. coli* metabolic reconstruction has served: As a content database where stoichiometrically and charge balanced reactions, and even pathways, have been incorporated into new reconstructions, As a database for defined metabolites, and as a source for a biomass objective function to query network content and functionality.

This influence has sparked an increase in the number of genome-scale network reconstructions that have been generated to formulate GEMs for a number of organisms. A detailed list of GEMs that have been developed, curated, and used for computation is given in Table 9.2. This table is a current snapshot of the

Fig. 9.4 Appearance of organism-specific genome-scale reconstructions and applications of the E. coli metabolism reconstruction. The genome-scale reconstructions for metabolic networks that have appeared every two years since the release of the first GEMs in 2000 (see Table 9.2) and the number of published studies the have appeared utilizing the *E. coli* GEM (Feist and Palsson 2008). Since the release of the first GEMs for *E. coli* (Edwards and Palsson 2000) and that of *Haemophilus influenzae* (Edwards and Palsson 1999), there has been a significant increase in both the number of genome-scale reconstructions and studies focused on the *E. coli* GEM for every time period

available reconstructions and a continually updated version can be found online (http://systemsbiology.ucsd.edu/*In Silico* Organisms/Other Organisms). Additionally, Fig. 9.4 shows the number of genome-scale reconstructions that have been developed over two year periods (for the reconstructions listed in Table 9.2). The number of reconstructions generated for each period has increased since the release of the first genome-scale reconstructions for *Haemophilus influenzae* in 1999 (Edwards and Palsson 1999) and *E. coli* in 2000 (Edwards and Palsson 2000). Furthermore, the number of published studies utilizing the *E. coli* GEM has also increased significantly over time resulting in the applications outlined in the sections below (Feist and Palsson 2008).

Modeling strategy and philosophy: Models are a formal way of accounting for our knowledge about the phenomena being described. When describing biochemical reaction networks formally, we need to deal with the 'links' (i.e., the reactions) between 'nodes' (i.e., the compounds). Our knowledge about links between biological molecules varies; from the abstract to the specific (Fig. 9.5). Statistical models are built on correlations and a black box approach that is not mechanism based. Specific mechanism-based models are based on knowledge of chemistry, kinetics, and thermodynamics. Given the fact that kinetic and thermodynamic information is hard to obtain on a large-scale, stoichiometric models stop one step short of full specification (in the spectrum conveyed in Fig. 9.5). The result is that we have chemistry (and its genetic basis) and network structure used as the foundation for building a mathematical description of network functions. Such models do not have a unique solution

Fig. 9.5 The different levels of knowledge used to generate biological models. Our knowledge about links in biochemical networks varies. At one extreme, the information is abstract and often takes the form of *black-box* correlations. At the other, we have detailed chemical mechanisms with kinetic and thermodynamic information. Stoichiometric models would be second from the *right*, accounting for mechanisms, but not incorporating kinetic and thermodynamic information

(e.g., see (Palsson 2006) and below). The lack of kinetic information can be dealt with by: (1) examining the properties of the entire set of solutions (i.e., the solutions space) or (2) by using constraint-based optimization to find specific solutions in the space (Price et al. 2004a). The latter can be successful if we know the prevailing selection pressure on an organism. The combination of a network reconstruction that is based on a knowledge-base at the genome-scale and the inherent optimality properties of the selection process underlie the success of COBRA for a number of applications.

Constraint-based modeling methods: Over the past quarter century, there has been a growing number of computational tools developed to interrogate biological networks and models (Breitling et al. 2008, Palsson 2006, Price et al. 2004a). Owing to its early development, the *E. coli* reconstruction and model has been a popular target for initial screening and development of a number of these methods. In this section, we introduce basic concepts common to most of these methods and describe in more detail those methods that were used in the studies presented in this chapter. The interested reader is encouraged to refer to recently published reviews presenting the constraint-based modeling methods in more detail (Breitling et al. 2008, Palsson 2006, Price et al. 2004a).

Mathematical description of the reconstruction: The metabolic reconstruction consists of a list of biochemical transformations known to take place in the target organism. This reaction list can be readily converted into a mathematical, computable format by using any available parser (e.g. in COBRA toolbox (Becker et al. 2007)). Using a parser, the stoichiometric coefficients are extracted for the individual reactions and entered in the cell of the stoichiometric matrix, also called the S matrix (Fig. 9.6). In this S matrix, every row corresponds to a metabolite and every column corresponds to a network reaction. Note that a typical S matrix

Fig. 9.6 The structure and application of constraints to networks. Shown are the components (Reaction network) and the engineering approaches and equations used to model a reconstructed network. The stoichiometric matrix is a mathematical representation of a reconstructed network and the steady-state assumption is used in a number of COBRA approaches, including flux balance analysis. The *bottom* of the diagram depicts how an unbound space can be confided to a solution space in which a network must behave by imposing the governing physiochemical constraints on a system (e.g., thermodynamic constraints)

is very sparse (< 1% non-zero entries) as many biochemical transformations are bi-linear, and the majority of metabolites appear only in few metabolic reactions. Only a few metabolites, such as protons, water, and ATP, are highly connected in a metabolic network, and participate in many metabolic reactions. Many studies have concentrated on studying the topological features of metabolic networks and the S matrix (see Section 9.5.4 or (Feist and Palsson 2008)).

The multiplication of this S matrix with a flux vector ν , containing flux values for all reactions v_j in S, results in a vector listing the changes in concentrations of all metabolites x_i over time:

$$
S \bullet \nu = \frac{dx}{dt} \tag{9.1}
$$

The constraint–based modeling approaches are based on the steady state assumption (Fig. 9.6), which assumes that the change of metabolite concentration over time is zero:

9 Genome-Scale Reconstruction, Modeling, and Simulation 163

$$
S \bullet v = \frac{dx}{dt} = 0 \tag{9.2}
$$

This assumption is valid for the metabolic reactions as the time scale of the reaction rates is much smaller (milliseconds range) than the doubling time of a cell, which is on the order of hours. Due to this time-scale separation, the metabolic network is essentially in a steady state during cell replication, and as a consequence, intracellular metabolites are not allowed to accumulate. This restriction, imposed by Equation (9.2), is known as the mass-balance constraint (Fig. 9.6).

Further constraints may be added to the reconstruction, leading to the conversion of the reconstruction to a condition-specific model. Such constraints can include thermodynamic (i.e., reaction reversibility), regulatory (e.g., expression of an enzyme), topological (i.e., composition and connectivity of network), and environmental (e.g., presence/absence of a specific carbon source).

Interrogation of the steady state solution space: In most cases, the set of equations encoded in the S matrix are underdetermined, meaning that there are more variables (fluxes v_j for $j = 1...n$) than there are equations (mass-balances for each metabolite x_i for $i = 1...m$. As a consequence, there is no single solution or flux vector v satisfying all the equations, but rather there are many possible flux vectors. This set of possible flux vectors is called the steady-state solution space. Each flux vector v , satisfying the given model constraints, is called a functional state of the network. This term functional state can be seen as analogous to the traffic pattern of the road mesh in a large city. The road mesh would correspond to the metabolic network and the traffic pattern, which shows high traffic and low traffic on the highways, corresponds to the functional state of the road system. Clearly this traffic pattern will be very different in the afternoon during rush hour versus the traffic pattern found late into the night. This example highlights the idea that one network can have many distinct functional states.

Functional states of a network can be determined using different mathematical approaches. In the COBRA approach, there is a distinction between biased and unbiased methods. Biased methods require the statement of an objective function, such as a biomass formation reaction or a byproduct secretion reaction by the metabolic network. This objective function is then maximized (or minimized) to obtain a functional state leading to the maximal (or minimal) flux value of the objective function. In contrast, unbiased methods explore the entire steady state solution space by determining a representative subset of possible functional states that can be analyzed in a statistical manner. Examples of unbiased methods are uniform sampling (Almaas et al. 2004, Price et al. 2004b, Thiele et al. 2005a, Wiback et al. 2004) and extreme pathway analysis (Papin et al. 2002, Price et al. 2003).

In many COBRA applications, it is assumed that the aim of a living cell is to grow as fast as possible to outgrow competitors and thus to use available nutrients mainly for biomass production. Hence, many COBRA applications are used in conjunction with the maximization of the biomass production rate. For example, gene essentiality can be determined *in silico* where the essentiality of every gene is tested to see whether the metabolic network is still able to produce biomass despite the *in silico* disruption of a gene (see Fig. 9.9). Other examples in this chapter discuss metabolic engineering applications, where the metabolic network is modified in such way that it produces a desired byproduct while maintaining a certain biomass production capability. Many industrially-interesting byproducts are produced by cells when they cannot produce biomass (e.g., due to nitrogen or phosphate limitations). Thus, the byproduct and biomass production are competitive, or 'orthogonal' to each other. COBRA has been successfully used to couple the byproduct production with the biomass production by deleting certain metabolic genes, thereby redirecting carbon fluxes in the metabolic networks (see below). The byproduct coupling to biomass production forces the organism to produce the desired byproduct in order to obtain the cellular objective of biomass production.

9.4 Applications and Uses of the *E. coli* **Metabolic Reconstruction**

Ask not what you can do for a reconstruction, but what a reconstruction can do for you: The *E. coli* reconstruction and GEM has been adapted for a broad number of uses by research groups around the world. Studies utilizing the reconstructed *E. coli* network range from pragmatic to theoretical applications and address a wide range of questions. These uses can be further categorized into five areas which include: (1) metabolic engineering, (2) biological discovery, (3) assessment of phenotypic behavior, (4) biological network analysis, and (5) studies of bacterial evolution (Fig. 9.7). A more extensive review of these uses has recently appeared (Feist and Palsson 2008), as well as an additional review on metabolic engineering efforts with *E. coli* and other organisms (Kim et al. 2008). Here, key examples of uses of the *E. coli* reconstruction in each of these fields will be presented to demonstrate the utility of the reconstruction and modeling process.

Fig. 9.7 Spectrum of uses of the of the genome-scale E. coli metabolic network reconstruction. Uses of the *E. coli* metabolic reconstruction can be categorized into 5 different areas. Furthermore, these categories can be arranged in order of addressing more practical (e.g., generating a production strain) or more basic (e.g., understanding horizontal gene transfer) questions

9.4.1 Metabolic Engineering

Metabolic engineering efforts utilizing the GEM of *E. coli* have focused on exploring overproduction for a number of products. Three examples in which computation and experimental construction were used to achieve overproduction will be discussed here. The first two examples utilized the *E. coli* GEM to explore the

production of the amino acids L-valine (Park et al. 2007) and L-threonine (Lee et al. 2007) in *E. coli*, and each has demonstrated the broad usage of GEM-aided computation for strain design.

Production of L-threonine: In the first study, GEM-aided modeling was employed in three different areas to increase the production of L-threonine to industrial titers (Fig. 9.8) (Lee et al. 2007). In one instance, *in silico* modeling was used to identify the optimal activity of a key enzymatic reaction towards maximum Lthreonine production using a parametric sensitivity analysis that compared reaction activity to L-threonine production rate. The optimal activity prediction was subsequently used to tune the over-expression of the gene that encodes for this enzymatic reaction through comparison to base line activity, and the result was a production increase. This method proved to be vital to the success of this strain, as a previous transcription profiling guided attempt at over-expression resulted in an undesirable surplus of activity that was detrimental to L-threonine production.

For the same strain, a GEM-aided flux analysis in conjunction with mRNA expression data levels guided the elimination of negative regulation on a gene, which encoded for a reaction that channeled flux towards the final product. The third use of the GEM for the design of this strain occurred when an unwanted byproduct was observed in the culture medium and computation was utilized to divert the flux from this byproduct to L-threonine (Lee et al. 2007) through over-expression of another key gene encoded activity.

Fig. 9.8 Three different areas where modeling was incorporated to increase strain production. Areas of model-driven strain improvement utilized to overproduce L-threonine in *E. coli* (Lee et al. 2007). (**a**) Shown is a graph that provides the computed relationship between L-threonine production and the activity of particular reaction. This *in silico* parametric sensitivity analysis guided the level of expression necessary for increased production of the amino acid in the strain. (**b**) Given is a map of central metabolism representing the metabolic reconstruction of *E. coli*. In the analysis, expression data was mapped onto the network to guide the elimination of negative regulation and the network was used to overexpress a reaction that diverted flux away from a byproduct (byproduct elimination) towards the desired product

Production of lycopene: Lycopene is an important intermediate in the biosynthesis of many carotenoids, and it is used for food coloring as it possesses a strong color (bright red) and is non-toxic. To increase the production of an already highproducing strain, a systematic computational search was developed (Alper et al. 2005b) to explore the *E. coli* metabolic network and report gene deletions that diverted metabolic flux towards the desired product. This process resulted in a knockout strain that, when constructed, showed a two fold increase in the production of lycopene over the parental strain. In this analysis, the minimization of metabolic adjustment (MOMA) computational algorithm (Segre et al. 2002) and the IJE660 (Edwards and Palsson 2000) *E. coli* GEM were utilized to sequentially examine additive genetic deletions that would improve lycopene production while maintaining cell viability. It was found that this computational approach yielded a twofold increase in production rate over a previously engineered overproducing strain and an eightfold increase over wild-type production harboring only a lycopene biosynthesis plasmid (Alper et al. 2005b). In addition, the strain designs identified computationally were compared to mixed combinatorial transposon mutagenesis, and it was found that the maximum production observed could be designed solely using the systematic GEM-aided computational method (Alper et al. 2005a,b). Furthermore, a deleterious effect was observed when targets identified in individual computational designs were combined in an attempt to achieve an overall more desirable phenotype. Thus, the overall systematic effects from individual designs were not additive and needed to be interpreted in the context of the entire network.

Production of L-valine: This model-driven example of metabolic engineering demonstrates the use of applying a systematic computational search algorithm (Alper et al. 2005b) to the updated *E. coli* GEM MBEL979 (Lee et al. 2005) (similar to the *i*JR904 GEM (Reed et al. 2003)) to improve L-valine production. In this analysis, the *in silico* computation of beneficial knock-outs to divert flux towards the desired product once again resulted in a significant increase (greater than twofold) in the production of the desired metabolite over an existing overproducing strain (Park et al. 2007). A number of additional metabolic engineering approaches to increase overproduction were performed by, (i) relieving feedback inhibition and regulation through attenuation, (ii) removing competing pathways, (iii) up-regulation of primary biosynthetic pathways, and (iv) over-expression of export machinery. When compared to each of the other individual strain modifications, the *in silico* GEM aided interventions resulted in the greatest increase in L-valine production (Park et al. 2007). Taken together, this and the previous study demonstrate the broad applications for which GEMs can be utilized to design strains not only in a *de novo* fashion, but to make further improvements on strains through integrating and interpreting experimental data.

9.4.2 Biological Discovery

The GEM of *E. coli* can be used as a guide to discovery. There is still a significant amount of information missing relating to gene functions in *E. coli* (Riley et al. 2006), and the content contained in the *E. coli* reconstruction can be queried and analyzed to first, determine the current gaps in our knowledge of the organism and second, design experiments to specifically fill uncovered gaps in the knowledge landscape. Two examples of model-driven discovery are presented, and these studies should form the basis for further analysis. To uncover the genetic basis for experimentally observed functions in *E. coli*, the studies combined GEM-aided computation with guided experimentation.

Systems approach to refining genome annotation: The first study utilized an iterative process (Reed et al. 2006b) in which, (i) differences in modeling predictions and high-throughput growth phenotype data were identified, (ii) potential missing reactions that remedy these disagreements were algorithmically determined, (iii) bioinformatics was utilized to identify likely encoding ORFs, and (iv) resulting targeted ORFs were cloned and experimentally characterized. Application of this process led to the functional characterization of eight ORFs that are involved in transport, regulatory and metabolic functions in *E. coli* (Reed et al. 2006b). The discovery process was aided by a high-throughput growth phenotyping analysis and the genome-wide single-gene mutant collection (Baba et al. 2006), along with other characterization analyses such as targeted expression profiling. This work was the first such example of model-driven discovery of genome content aided by a metabolic network reconstruction.

Genetic basis of orphan reactions: The second GEM-based analysis that resulted in ORF discovery utilized network topology to examine orphan reactions in the *E. coli* network (i.e., reactions known to exist in *E. coli* that have not been linked to an encoding gene) identified by network topology-based gap-filling algorithms (Chen and Vitkup 2006, Kharchenko et al. 2006, 2004). The basic premise behind these algorithms is the utilization of an orphan reaction's network neighbors as constraints to assign metabolic function. With the resulting tentative ORF assignments, biochemical characterization studies utilizing genetic mutants (Baba et al. 2006), analysis of growth under different substrate conditions, and expression data were all utilized to characterize and assign function to an orphan ORF that is responsible for a metabolic conversion that has been known for 25 years (Fuhrer et al. 2007). These two studies are early examples of how GEM computation can lead to the discovery of new genetic and biochemical content in an organism.

9.4.3 Assessment of Phenotypic Behavior

Researchers have utilized the *E. coli* GEM to better understand the coordinated functions of the cell and observed physiological outcomes. Computations seeking to predict cellular phenotypes have been performed under a range of genetic and environmental conditions, and phenotypic assessment has received the most attention in terms of publication and tool development. Here, we outline computational tools developed to analyze the *E. coli* GEM in each of the two major areas of phenotypic assessment, studies of (i) network perturbation/essentiality, and (ii) the incorporation of thermodynamic information.

Fig. 9.9 Gene-deletion analyses utilizing the E. coli GEM. Analyses of gene essentiality in the *E. coli* metabolic network. (**a**) A table of results from an analysis performed using the *i*JE660 GEM of *E. coli* where experimental phenotypes were collected from bibliomic data. Results are scored as + or − meaning growth or no growth determined from *in vivo*/*in silico data*. The ± indicates that suppressor mutations have been observed that allow the mutant strain to grow. In 68 of 79 cases the *in silico* behavior is the same as the experimentally observed behavior. Each column represents a different carbon source. (**b**) This heat map characterizes the agreement between ORFs predicted to be essential using the *i*AF1260 GEM of *E. coli* (Feist et al. 2007) and those experimentally determined (Baba et al. 2006, Joyce et al. 2006). The enlarged region details how each row corresponds to a computationally predicted essential ORF (188 total). A dark row indicates the condition under which each ORF was found to be essential. For example, *folP* was predicted to be an essential ORF for the biosynthesis of folate in *i*AF1260 under these conditions, but was not identified as essential by Baba et al. (2006). The different columns show at which level each gene in the overall column was found to be essential on. With the advancement of both experimental data and model coverage, analyses of this type have reached the genomic scale

In silico **perturbations**: A set of distinct computational methods using GEMs has been developed to determine the physiological state of *E. coli* (and other cells for which a GEM exists) after genetic perturbations (Segre et al. 2002, Shlomi et al. 2005, Wunderlich and Mirny 2006). These methods were analyzed to examine the effectiveness of predictions when compared to experimental data (Fig. 9.9). Whereas comparisons to flux data from wild-type and *E. coli* mutants reveals that one of the computational algorithms, MOMA (Segre et al. 2002), provided better predictions for transient growth rates (early post perturbation state), another algorithm, ROOM (Shlomi et al. 2005) (and basic FBA), was found to be more successful in predicting final steady-state growth rates and overall lethality (Shlomi et al. 2005). These two algorithms have been utilized, in addition to basic FBA, for genome-wide essentiality screens. Aiding the effort is the recent availability of a comprehensive single-gene knock-out library for *E. coli* (Baba et al. 2006) which has been utilized for comparison with GEM computation (Feist et al. 2007, Joyce et al. 2006). Touching on the predictive capability of GEM computations, it was found that the *E. coli* GEM was able to predict the outcomes of adaptively evolved strains to a high degree (78%) when knock-out *E. coli* strains were grown in a number of different substrate environments by examining growth rates at the beginning and end of adaptive evolution (Fong and Palsson 2004). Genetic perturbations have played a key role in the study of the genotype-phenotype relationship in biology, and GEMs can be used to mechanistically interpret the results and predict the outcomes of such perturbations.

Adding thermodynamic information: The incorporation of thermodynamic information with GEMs is an effort that is progressing rapidly and should increase the predictive capabilities of genome-scale modeling through the addition of further governing physico-chemical constraints. Furthermore, the addition of thermodynamics enables the analysis of metabolomic data in the context of a reconstruction. A study utilizing high-throughput metabolomic data and GEMs resulted in the proposition of likely regulatory interactions by deciphering the metabolite concentrations in the context of overall network functionality (Kümmel et al. 2006). Not only did the metabolomic data benefit computations by constraining the system using physiological measurements, but the computational predictions were also able to validate quantitative metabolomic data sets for consistency through providing a functional context to relate metabolite concentrations. This application is one example of how metabolomic data will directly influence modeling. Metabolite concentration data is likely to greatly influence future metabolic modeling due to its intimate connection with GEM content.

9.4.4 Biological Network Analysis

Although there is still much to learn about the metabolism of *E. coli* and how a model-driven approach can be used to uncover these unknowns, the wealth of knowledge collected and represented in the current *E. coli* reconstruction makes it an ideal platform for network analyses. Researchers have been taking advantage of this fact and have centered network analyses on probing and uncovering the properties of biological networks in general. In this section, we discuss a key analysis based on the *E. coli* GEM and the implications drawn from such analyses.

One noteworthy study utilizing the *E. coli* network examined thousands of different potential growth conditions and resulted in the observation of a 'high-flux backbone' in *E. coli* that both, (i) carried high levels of flux across the different environmental conditions, and (ii) was composed of a relatively small set of enzymatic reactions (Almaas et al. 2004). This result can be of practical importance for synthetic biology efforts aimed towards manipulating flux within biological systems. Furthermore, this finding was hypothesized to be a universal feature of metabolic activity in all cells and was consistent with flux measurements from 13C labeling experiments (Almaas et al. 2004).

Overall, studies of network analyses have a common systems biology theme: the development and subsequent demonstration of methods that identify sets of reactions or metabolites with correlated or coordinated functions and systematic relationships. The systems biology that these methods enable and demonstrate has the potential to influence the more practical applications already outlined. The role that the *E. coli* GEM has taken is a comprehensive and curated set of up-to-date metabolic knowledge that provides a scaffold for large-scale computations.

9.4.5 Studies of Bacterial Evolution

The GEMs of *E. coli* have been used to examine the process of bacterial evolution (Pal et al. 2005a,b, 2006). Specifically, the network reconstructions have been used to interpret adaptive evolution events (Pal et al. 2005a), horizontal gene transfer (Pal et al. 2005a,b) and evolution to minimal metabolic networks (Pal et al. 2006). These studies, which utilize the *E. coli* reconstruction as an organism-specific genetic and metabolic content database and the corresponding GEM have been able to provide insight into evolutionary events through combining known physiological data (e.g., in various environmental conditions) with hypotheses and *in silico* computation. Examination of the evolution of minimal metabolic networks through simulation demonstrated that it was possible to predict the gene content of close relatives of *E. coli* by examining the necessity of genes and reactions in the overall context of the system functionality for a specific lifestyle (Pal et al. 2006). Similarly, by re-examining network functionality in a number of different environments, and through the utilization of comparative genomics, it was shown that recent evolutionary events (i.e., horizontal gene transfer) likely resulted from a response to a change in environment (Pal et al. 2005a). Furthermore, computational analysis led to the additional conclusion that these horizontal gene transfer events are more likely when the host organism contains an enzyme that catalyzes a coupled metabolic flux related to the transferred enzyme's function (Pal et al. 2005a,b). Taken together, these studies demonstrate the importance of having high-quality curated reconstructions to enable studies on an organism's response to environmental changes and on the fundamental forces driving bacterial evolution.

9.5 Need for New *In Silico* **Methods and Applications**

We now know how to represent BiGG data in either a stoichiometric format or in the form of causal relationships (Gianchandani et al. 2006) and how to use this data to perform several lines of computational inquiries. Computational query tools of GEMs will continue to be developed. New advances in these query tools will likely include, (i) modularization methods, (ii) use of fluxomic data, and (iii) eventually, kinetic information.

Modularization: As the scope and content of the reconstruction grows, the need to modularize its content becomes more pressing. Fine or coarse-grained views of cellular processes are needed for different applications.

Fluxomics: Currently, computational limitations force the reduction in network size for the analysis of isotopomer data. Given the systemic nature of fluxomic data and its phenotypic relevance, there is a pressing need to increase the size of the networks that can be utilized for experimental measurement and estimation of flux states. A network reconstruction will both guide the content that is needed for analyzing fluxomic data and offer a starting point for a rational reduction to generate relevant models in the meantime.

Kinetics/thermodynamics: Although detailed kinetic models of microbial functions may currently be mostly of academic interest, they will most likely be able to be constructed in the mid-term based on advances with metabolomic and fluxomic data, in addition to the developments that are occurring with the incorporation of thermodynamic information. Such large-scale kinetic models are likely to differ from those resulting from traditional approaches for construction of kinetic models as they come with different challenges.

9.6 Closing

The process underlying the *E. coli* metabolic reconstruction has pioneered many approaches, methods, and studies in the systems biology of microbial metabolism. This effort has effectively put a mechanistic basis into the genotype-phenotype relationship. In fact, this relationship is now broken down into four steps:

- (1) Components (a large knowledge base, BiGG), leading to networks (the reconstruction process resulting GENRE), leading to *In silico* Models (GEMs), leading to Phenotypic States (estimated by COBRA methods).
- (2) GEMs will allow for gap-filling and systematic biological discovery (Breitling et al. 2008) and for understanding of complex biological processes (see Chapter 15).

Predictive models also allow for experimental strain design. In fact, in engineering, there is '*nothing more practical than a good theory*.' As this chapter demonstrated, genomics and high-throughput technologies have enabled the construction of predictive computational models. The scope of such predictions is limited at the moment, but with the growing scope and coverage of genome-scale reconstructions and advancements in the development of computational tools, this scope will broaden. Not only will GEMs influence design in synthetic biology, but also their help with discovering cellular content will provide a more complete picture of the intra-cellular environment in which future synthetically engineered constructs and circuits will be placed. The impact of GEMs on synthetic biology is thus likely to be notable, ranging from the provision of the cellular context of a small-scale gene circuit design to engineering of the entire genome-scale network towards fundamentally new and useful (i.e., production) phenotypes.

Finally, we can speculate about the deep scientific impact that comprehensive predictive GEMs will have on our understanding of the living process. A comprehensive view of cellular functions will allow us to study the fundamental properties of both the underlying energy and information flows in living organisms. Such a view is likely to deeply affect our understanding of both distal and proximal causation in biology.

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